

Sealed operation, and circulation and purification of gas in the HARPO TPC

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December 11, 2015

Abstract

HARPO is a time projection chamber (TPC) demonstrator of a gamma-ray telescope and polarimeter in the MeV-GeV range, for a future space mission. We present the evolution of the TPC performance over a five month sealed-mode operation, by the analysis of cosmic-ray data, followed by the fast and complete recovery of the initial gas properties using a lightweight gas circulation and purification system.

1 Introduction

High-energy astronomy is hindered by the sensitivity gap between the energy ranges over which the Compton and the pair telescopes are efficient, making the γ -ray sky in the range 0.1 – 100 MeV almost unexplored. Furthermore no polarimeter with a decent sensitivity above 1 MeV has ever been flown in space.

On the low-energy side of the gap, a number of projects are aiming at an improved sensitivity with respect to COMPTEL. Of these, those using stacks of silicon wafers without any tungsten converter can also

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detect pair conversions [1–3] with an angular resolution of $\approx 1.5^\circ$ @ 100 MeV, a significant improvement over that of the Fermi/LAT ($\approx 6^\circ$ @ 100 MeV [4]). These stacks can have some sensitivity to polarisation if the wafers are thin enough and provided that matching can be performed in the first wafer after the conversion wafer, i.e. that each of the hits in the x direction can be assigned to one of the two hits in the orthogonal (y) direction. In subsequent wafers, the azimuthal information carried by the pair is washed out by multiple scattering [5].

In the quest for even thinner wafers, it is tempting to use an even less dense detector such as a gas detector. In the case of an optimal tracking in the presence of multiple scattering such as with a Kalman filter, an even better angular resolution of $\approx 0.4^\circ$ @ 100 MeV can be achieved [6] providing a point-like source sensitivity as low as $10^{-6}\text{MeV}/(\text{cm}^2\text{s})$ @ 100 MeV with a 10 kg gas time projection chamber (TPC), while that of COMPTEL plateaued at $\approx 10^{-4}\text{MeV}/(\text{cm}^2\text{s})$ [7]. In such gas detectors the dilution of the azimuthal information carried by the pair is close to unity, so that the sensitivity to polarisation is excellent [8]. Finally and most importantly track matching can easily be performed by comparing the distribution of the energy deposited along the tracks in the x and y directions thanks to the large fluctuation in the local deposition of energy along the track [9]. These unique features come at the cost of the need to maintain a good-quality gas in which the collection of the ionizing electrons is not affected by impurities with a large electron attachment. For example the EGRET gas (99.5% neon, 0.25% argon, and 0.25% ethane) became contaminated by the breakdown of ethane by spark chamber firings and to a small degree by residual outgassing. The five planned gas exchanges were carried out once per year, after which the lack of fresh gas became a serious issue [10]. These concerns have triggered the development of detectors built out of low outgassing rate materials. For example, for the GEMS X-ray polarimeter TPC, the design of which involves copper clad liquid crystal polymer (LCP) gas electron multipliers (GEM), polyetheretherkeytone (PEEK) dielectric structure on a gold-plated titanium frame, a 23 year lifetime is expected [11].

We report here on a gas purity study performed with the HARPO detector, a TPC prototype of a $\gamma \rightarrow e^+e^-$ telescope built with techniques much better suited to our actual budget and that we have studied in the laboratory with radioactive sources [12] and with cosmic rays [13], and that we have tested recently [14] in a γ -ray beam.

2 Description of the HARPO TPC

HARPO is a 30 cm cubic TPC surrounded by 6 scintillator plates, each equipped with two wavelength shifter bars and two photomultipliers (PMTs) for trigger and background rejection. The gas container is an

aluminum cylinder designed to be operated from few mbar up to 5 bar in sealed mode (Fig. 1). With a mixture of 95% of argon 5.0 and 5% of iC_4H_{10} 3.5 and at a 220 V/cm drift electric field, the electron drift velocity is about 3 cm/ μ s. The TPC end-plate is a "hybrid" MicroPattern Gas Detector (MPGD) composed of two GEM foils [15] with 2 mm spacing above a 128 μ m amplification gap bulk-micromegas [16]. The anode collection plane is segmented in 2×288 strips in the two orthogonal x, y directions and the signal is read by an electronic chain composed of two T2K/TPC Front-End cards [17], two FEMINOS back-end cards and a Trigger Clock Module [18]. A simple coincidence between scintillators is used here for triggering on cosmic rays.

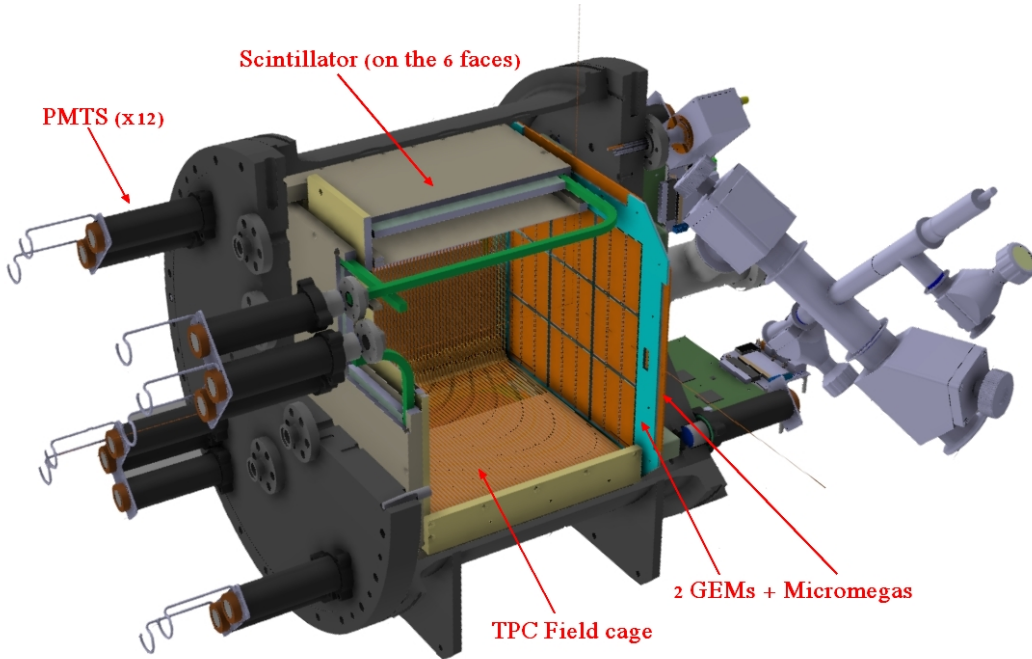


Figure 1: A 3D CAD drawing of the HARPO demonstrator.

3 Description of the circulation and purification system (CAPS)

We used a circulator composed of a simple turbine (Fig. 2, left). The gas was lead through an Oxisorb purification cartridge that can absorb up to 0.5 L H_2O and 0.1 L O_2 , with output purity as low as < 5 ppb for O_2 and 30 ppb for H_2O [19]. We measured the gas flow with a F-111B Bronkhorst mass flowmeter (Fig. 3, I10) that was calibrated for our gas mix at 2 bar and 20 $^{\circ}C$. We monitored the gas pressure

and temperature with a Keller EV-120/PA(A)-33X sensor (Fig. 3, I4 and I5) with ranges of 0 – 50 °C and 0 – 10 bar and resolutions of 0.01 °C and 1 mbar. The outgassing areas inside the vessel amount to 0.82 m² Aluminum, 3.50 m² PVC, 0.92 m² PCB, 1.63 m² PVT, for a gas volume $V_{\text{det}} = 84$ L.

We tested the gas circulation system with a helium leak detector (ASM 310) down to $3.3 \cdot 10^{-8}$ mbar · L/s, after which we opened the connection between the CAPS and the detector. This caused a pressure drop from 2015 mbar to 1960 mbar at 27 °C, with a flow of 1.5 L/h. On July 20 2015, we opened the Oxisorb purificator in vacuum which lead to a new pressure drop to 1932 mbar when the connection to the detector was opened. The CAPS then ran continuously, with a flow through the filter of $f_{\text{circ}} = 1.4$ L/h. In these conditions we expect the water and oxygen content in the gas, C , to evolve following a law of the form:

$$C = C_0 e^{-t/\tau_{\text{purif}}} + K_{\text{cont}} \tau_{\text{purif}}, \quad (1)$$

with $\tau_{\text{purif}} = V_{\text{det}}/\epsilon_{\text{purif}}f_{\text{circ}}$, where C_0 is the original content and K_{cont} is the contamination rate from leaks and outgassing. In case the purification efficiency of the filter is $\epsilon_{\text{purif}} = 1$, the purification rate is equal to $\tau_{\text{purif}} = 60$ h.

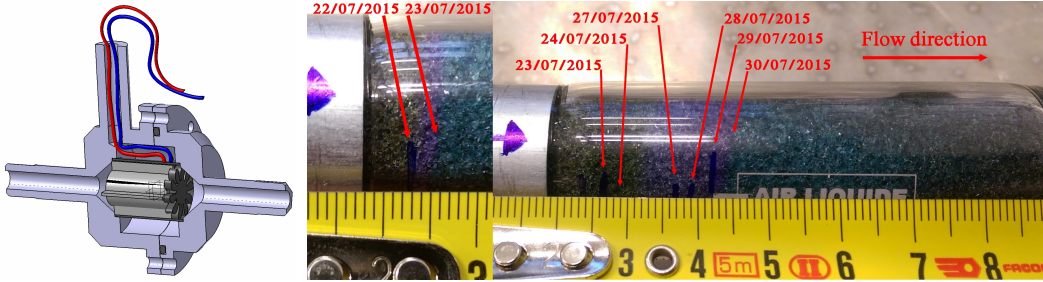


Figure 2: (Left) Sectional view of the circulator (CEA patent BD15730SG). (Right) Picture of the Oxisorb cartridge after 10 days of continuous use.

The Oxisorb purificator color ranges from blue to dark brown with the quantity of H₂O and O₂ absorbed. At the beginning of the purification, we observed a purple ring of 3 mm thickness growing 3 mm per day during the first 4 days. It stabilised at 12 mm after a few days (Fig. 2). This indicates that the cartridge was never saturated.

Several gas samples were taken for analysis, which lead to density drops (lines 1, 2, and 3 on Fig. 7). Before and after every operation that affected the gas in the vessel, data was taken with the TPC for consistency checks.

4 Data taking and analysis

We took cosmic-ray data once per week from February to July, with the detector set so that the drift field was vertical. Most runs were one hour long, recording a few 10,000 events. After starting the purification on July 20 we took data more frequently. The high voltage was only turned on when taking data. Due to the vertical detector orientation, most of the recorded tracks are approximately aligned with the drift direction. Figure 4 shows the x, t and y, t projections for a typical event. We select events with a single track in each projection, with a high momentum muon selection based on the χ^2 cut of a linear fit. For each time bin t_i ($30 \text{ ns} \approx 1 \text{ mm}$), we obtain the charge deposited by a track, $Q(t_i)$, summing over all the channels associated with the track. After an angle-effect correction using the direction \vec{u} of the track,

$$\frac{dE}{dx}(t_i) = Q(t_i) \times u_z, \quad (2)$$

we obtain the $dE/dx(t_i)$ in ADC units, that is up to an overall factor that includes the gain of the amplification of both the MPGDs and that of the readout electronics. The resulting distribution (Fig. 5) is also affected by threshold and saturation effects. To mitigate them, we estimate the most probable value (MPV, solid line in Fig. 5) of the charge for each time bin with a Landau-function fit. From this distribution we extract :

The drift velocity. As the tracks are selected to traverse the full TPC length, the drift-time duration provides a measurement of the drift velocity.

The gain. The height of the distribution at short drift times is proportional to the amplification gain.

The attenuation. The signal loss along the drift from electron capture is visible in Fig. 5. We fit the central part of the distribution, which is less affected by the threshold and saturation effects, with an exponential function.

These measurements are affected by systematic effects. Since the detector configuration was kept unchanged throughout the data taking, they cancel in the relative value compared to a reference run, chosen to be run 2008 (Feb. 18th), i.e. right after the fresh gas fill. Figure 6 shows the relative charge distribution as a function of the drift distance for several runs taken at a few weeks interval.

5 Results

For the first five months, while the TPC was sealed, we observe a steady degradation of the gain, of the drift velocity and of the electron attenuation (Fig. 7). After we turned the CAPS on, the drift velocity and

Table 1: Gas content from mass spectrometry (volume/volume). Note that the set-up was not sensitive to H_2O .

Sample #		1	2	3
Compound		July 08	Aug. 27	Sept. 17
iC ₄ H ₁₀	%	5.10	4.42	4.49
O ₂	ppm	180	<20	<20
CO	ppm	190	250	130
CO ₂	ppm	120	160	130
N ₂	ppm	620	890	850

the attenuation got back to their initial values. We corrected for the gain changes due to the density decrease induced by our extraction of gas samples and extension of the volume with the CAPS circuit, by adjusting the MPGD voltages. The pressure change also affected the drift velocity but to a negligible extent as shown by Garfield simulations [20], compared to the observed effect.

Gas samples were collected before (#1, line 3 on Fig.7), and respectively five weeks (#2) and eight weeks (#3) after activating the purification system. They were analysed in a high resolution ($R = 2800$) direct-injection THERMO MAT 271 mass spectrometer (Fig. 8, Table 1). The relative amount of O₂ and N₂ before purification is consistent with that of air, due to either leaks or outgassing. Contaminations by CO and CO₂ probably originate from outgassing from plastic elements.

After purification, the residual oxygen content is found to be lower than the detection threshold of the spectrometer. We also note a small but significant decrease of the amount of isobutane content upon purification after which it stabilized, an effect that is not understood.

6 Discussion

Garfield++ simulations [20] show that the degradation of the attenuation during sealed operation, attributed to electron attachment, is proportional to the oxygen content. After purification was started the effect disappears exponentially with a characteristic time $\tau_{\text{purif}} = 105 \pm 4 \text{ h}$ (Fig. 7). The observed time constant is larger than the computed value, most likely due to the limitation of gas circulation inside the almost closed field cage. We also observe a decrease of the drift velocity, which cannot be associated with the oxygen contamination and which is recovered after purification, and therefore that can be attributed to water contamination.

After purification, the drift velocity was measured to be $6.0 \pm 0.5\%$ higher than its initial value. From the Garfield simulation, this can be explained by the cumulative effects of the reduced gas pressure from 2.1 to 1.9 bar (+4%) and of the measured decrease of the isobutane content from 5.1% to 4.5% (+3%).

7 Conclusion

We have successfully operated a sealed TPC over five months, with a clear but manageable gas degradation. We used the data from cosmic rays in the TPC to monitor the relevant parameters of the gas (drift velocity, gain, attenuation). We then purified the gas with a simple, low-power circulation system and observed a fast and complete recovery of the initial parameters. If operated routinely, for example on a space mission, the system should therefore enable a stable performance for several years.

This work is funded by the French National Research Agency (ANR-13-BS05-0002)

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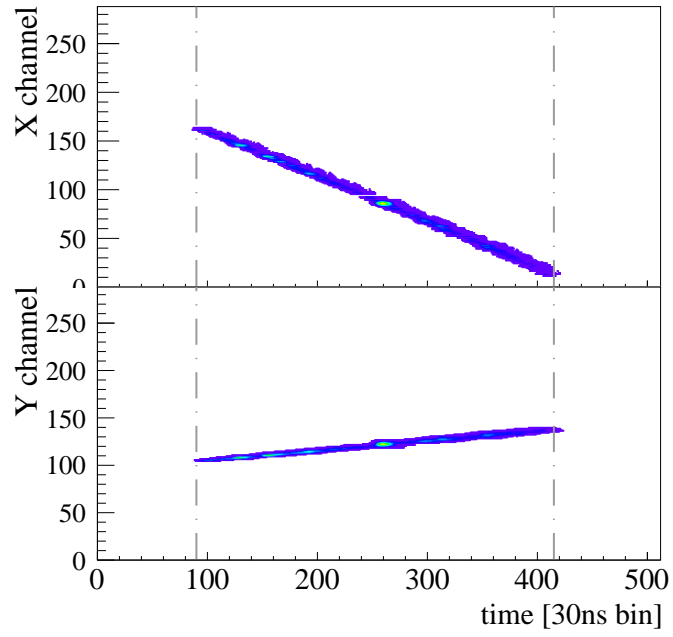


Figure 4: A cosmic-ray event in HARPO. The “maps” show the charge distributions in the projection X-time and Y-time.

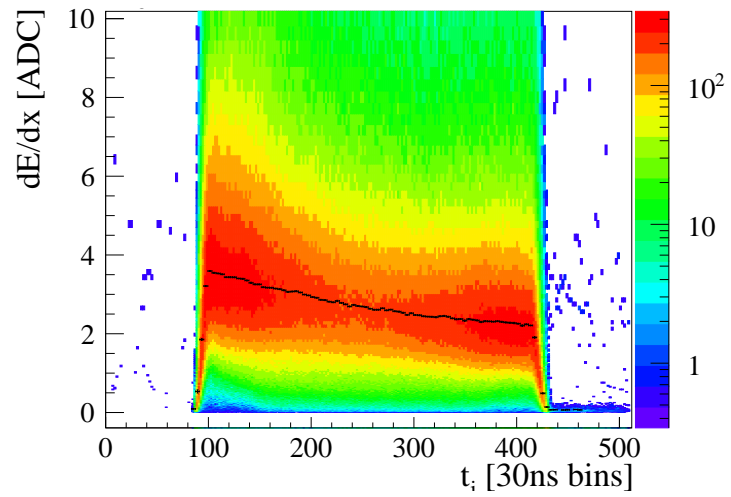


Figure 5: Normalised charge as a function of the drift time for a 6000 s cosmic-ray run.

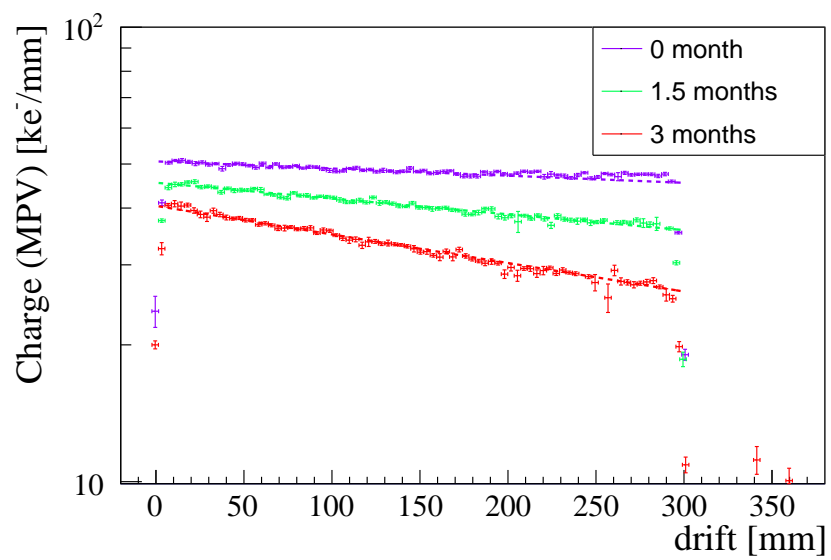


Figure 6: Charge as a function of the drift distance, for three runs, 6 weeks apart, normalised to the reference run 2008.

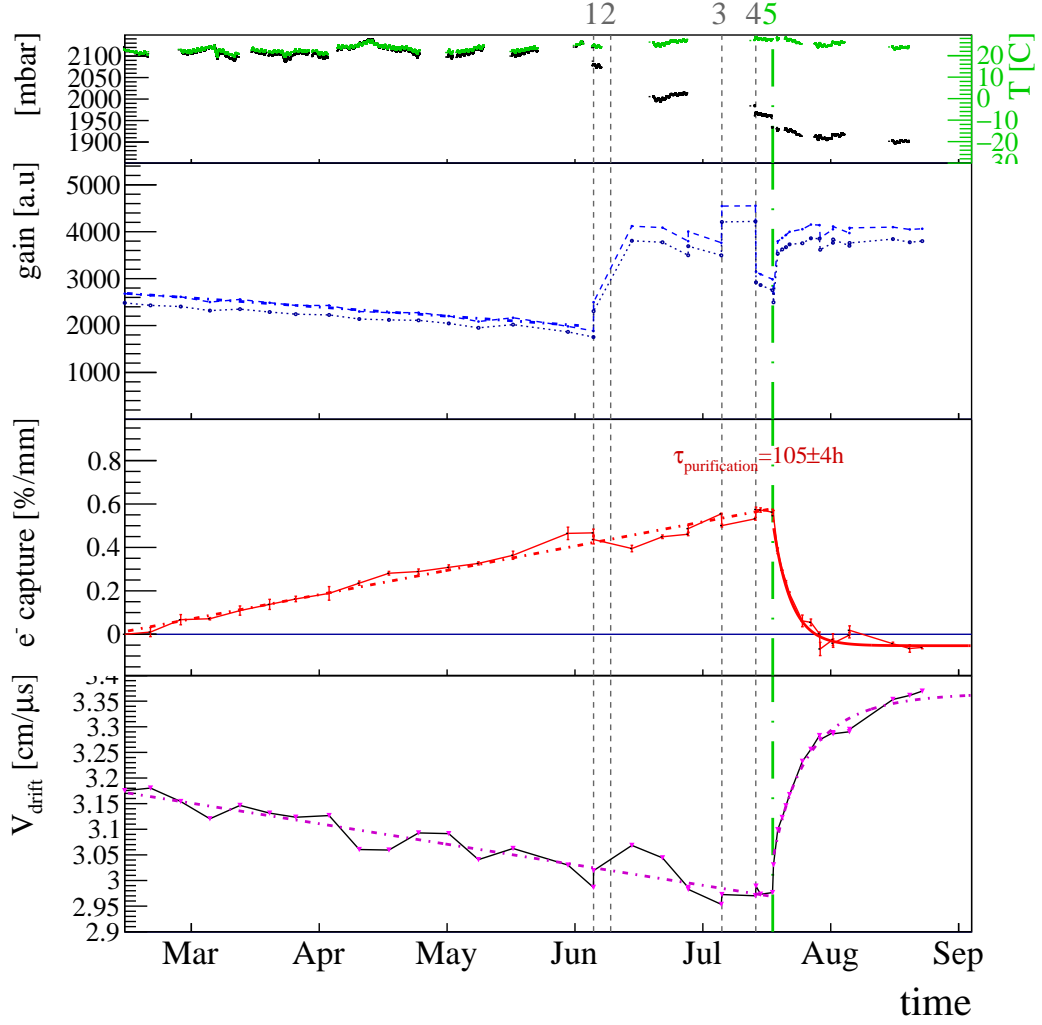


Figure 7: Time evolution of the gain, of the attenuation and of the drift velocity. We can see several variations of the gain due to the pressure loss when gas samples were extracted. For the first five months during which the TPC was sealed we see a regular degradation of all three parameters. On July 20 (line 5), the CAPS was turned on.

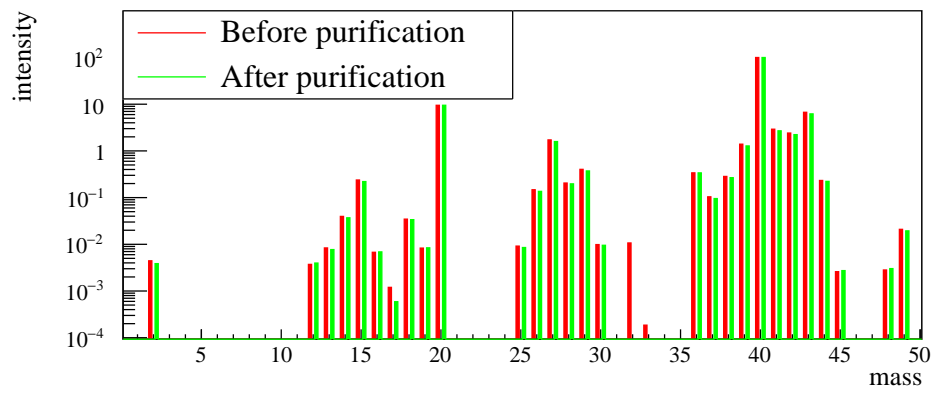


Figure 8: Mass spectroscopy results for samples taken before and after purification. The overall composition is similar, except for $N=32$ (O_2), that disappeared in the second sample.